

Rev 1: September 2018 FSN Ref: 446776

FSCA Ref: 446775

Date: 24.04.2024

Urgent Field Safety Notice

VARIOUS MAPLESON F ANAESTHETIC BREATHING SYSTEMS CONTAINING 0.5L RESERVOIR BAGS WITH OPEN TAILS

For Attention of*: MDSO's, All clinical staff, Managers and users of the above products

Contact details of local representative (name, e-mail, telephone, address etc.)* Chris Randall Wokingham Site Quality Manager Intersurgical Ltd. Crane house Molly Millars Lane Wokingham Berkshire RG41 2RZ Email: priority@intersurgical.co.uk Tel. No: 0118 9656 300 Fax: 0118 9656 356



Urgent Field Safety Notice (FSN)

VARIOUS MAPLESON F ANAESTHETIC BREATHING SYSTEMS CONTAINING 0.5L RESERVOIR BAGS WITH OPEN TAILS

Risk addressed by FSN

1. Information on Affected Devices* 1. Device Type(s)*			
Various Mapleson F Anaesthetic Breathing Systems			
 Commercial name(s) Mapleson F infant T-piece breathing system with 0.5L open tail bag, ≥ 1.8m Mapleson F Jackson Rees modification T-piece breathing system with 0.5L open tail bag, ≥ 1.8m 			
Paediatric, Mapleson F Jackson Rees modification T-piece with 0.5L open tail bag, and swivel elbow, ≥ 1.8m Paediatric, Mapleson F Jackson Rees modification T-piece with 0.5L open tail bag, and swivel elbow, ≥ 4.8m Paediatric, Mapleson F Jackson Rees modification T-piece with 0.5L open tail bag, and elbow, ≥ 2.8m Paediatric, Mapleson F Jackson Rees modification T-piece with 0.5L open tail bag, and elbow, ≥ 1.8m			
Paediatric, Mapleson F Jackson Rees modification T-piece with 0.5L open tail bag, and swivel elbow, ≥ 3.6m Paediatric, Mapleson F Jackson Rees modification T-piece with 0.5L open tail bag, and luer elbow, ≥ 3.6m Map/F 0.5L Open T/B Luer/Elb >= 2.4m Map/F 0.5L Open T/B Luer/Elb M/Line >= 1.8m			
Map/F 0.5L Open T/B Luer/Elb >= 1.6m Paediatric, Mapleson F Jackson Rees modification T-piece with 0.5L open tail bag, and luer elbow, ≥ 10.8m			
3. Unique Device Identifier(s) (UDI-DI)			
50302670622495030267062270503026706236250302670624305030267103164503026706225650302670622875030267062379503026706250850302671498105030267062263503026706234850302670623935030267062539			
4. Primary clinical purpose of device(s)*			
To deliver and remove anaesthetic and respiratory gases to and from a paediatric patient via a breathing system comprised of tubing and connectors and 0.5 L reservoir bag.			
5. Device Model/Catalogue/part number(s)*			
2120000 2121004 2121014 2121035 2121048			
2121000 2121005 2121019 2121042 2121053 2121002 2121011 2121024 2121045			
6. Software version			
N/A			
7. Affected serial or lot number range Any of the above with an expiry date from April 2024 to March 2029.			
8. Associated devices			
N/A.			



	2. Reason for Field Safety Corrective Action (FSCA)*	
2.	1. Description of the product problem*	
2.	Some devices contain a reservoir bag with a closed tail, when they should have a reservoir bag with open tail.	
	Correct – Open Tail Reservoir Bag Incorrect – Closed Tail Reservoir Bag	
2.	2. Hazard giving rise to the FSCA* If the incorrect closed tail reservoir bag is not identified during the routine pre use check as described in the product instruction for use, it could result in over pressurisation of the system leading to potential barotrauma.	
2.	3. Probability of problem arising	
	100% in the affected range.	
2.	4. Predicted risk to patient/users	
<u></u>	The risks associated with the identified fault have been reviewed, and whilst the probability of occurrence is low, we believe it is essential to address the issue promptly to further reduce the risk of any potential patient harm.	
2.	5. Further information to help characterise the problem	
	N/A	
2.	6. Background on Issue	
	Following a customer report from the market and subsequent thorough inspection and analysis of internal stock, we have identified a potential safety concern related to various Mapleson F paediatric anaesthetic breathing systems as listed above. Unfortunately some products have been manufactured with a 0.5 L reservoir bag with a closed tail which could result in over pressurisation of the system.	
2.	7. Other information relevant to FSCA	
	N/A	
	· · · ·	



	3. Type of Action to mitigate the risk*			
3.	1. Action To Be Taken by the User*			
	⊠ Identify Device □ Q Device	uarantine Device	C Return Device	e 🗆 Destroy
	□ On-site device modifica	tion/inspection		
	Follow patient management recommendations			
	☑ Take note of amendment/reinforcement of Instructions For Use (IFU)			e (IFU)
	⊠ Other □ No	one		
	Please distribute this Field Safety Notice to all potential users of the Mapleson F paediatric anaesthetic breathing systems listed above, within your facility. This is for their awareness of the potential problem and to carry out the following actions.			
	To ensure the safety of patien	ts we recommend t	he following action	S.
	1. Identify any potentially affeo above.	cted products from	he affected codes	and lot numbers listed
	2. All users must perform a thorough visual inspection and functional test before use of the products and lot numbers listed above, to confirm a patent gas pathway exists through the open tail of the reservoir bag to avoid over pressurisation of the system.3. Retain any affected sample(s) identified, and please report to us immediately.			
	Please note: This is not a product removal.			
	Please complete and return the Reply Form provided to priority@intersurgical.co.uk, to confirm receipt of this notice and that the necessary actions are being taken.			
	Please continue to report to Intersurgical any adverse events involving this product.			ng this product.
3.	2. By when should the action be completed?	FSN should be or	· · · ·	and awareness of this ntially affected stock
3.	3. Particular considerations f	or: N/A		
	Is follow-up of patients or i	review of patients' p	previous results rec	ommended?
	Not applicable.			
3.	4. Is customer Reply Require (If yes, form attached specifying			es
3.	5. Action Being Taken by t	he Manufacturer	I	
	Product Removal	□ On-site device	modification/inspe	ction
	□ Software upgrade	☑ IFU or labelling	•	
	⊠ Other	□ None	, 0-	



We have implemented corrective actions in manufacturing process to eliminate this problem for future supply. We will also be introducing a new instruction for use which will include the following Pre-Use Check in line with the recommended action 2. above:

If the product is supplied without an APL valve, the pressure within the system is controlled by the clinician through manipulation of the open tail on the reservoir bag. Check that a patent gas pathway exists through the open tail of the reservoir bag.

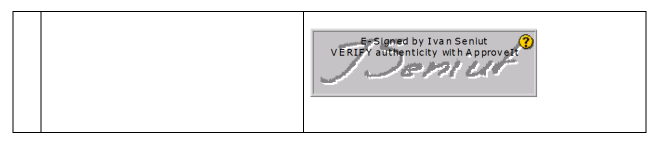
3	6.	By when should the action be completed?	One month from receipt of the FSN	
3.	7.	Is the FSN required to be c /lay user?	communicated to the patient	No
3	 If yes, has manufacturer provided additional information suitable for the patient/lay user in a patient/lay or non-professional user information letter/sheet? N/A 			

	4. General Information*		
4.	1. FSN Type*	New – Advisory Notice	
4.	2. For updated FSN, reference number and date of previous FSN	N/A	
4.	3. For Updated FSN, key new informa	ition as follows:	
	N/A		
4.	 Further advice or information already expected in follow-up FSN? * 	No	
	5. If follow-up FSN expected, what is	the further advice expected to relate to:	
4	N/A		
	6. Anticipated timescale for follow-up	N/A	
4	FSN		
4.	7. Manufacturer information		
	(For contact details of local representation		
	a.Company Name	Intersurgical Ltd.	
	b. Address	Crane House, Molly Millars Lane, Wokingham, Berkshire, RG41 2RZ	
	c.Website address	https://www.intersurgical.com/	
4.		ority of your country has been informed about this	
	communication to customers. *		
4.	9. List of attachments/appendices:	Customer Reply Form	
4.	10. Name/Signature	Ivan Seniut, Group Quality and Regulatory Affairs Director, Intersurgical	



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Transmission of this Field Safety Notice
This notice needs to be passed on all those who need to be aware within your organisation or to any organisation where the potentially affected devices have been transferred. (As appropriate)
Please transfer this notice to other organisations on which this action has an impact. (As appropriate)
Please maintain awareness on this notice and resulting action for an appropriate period to ensure effectiveness of the corrective action.
Please report all device-related incidents to the manufacturer, distributor or local representative, and the national Competent Authority if appropriate, as this provides important feedback.

Note: Fields indicated by * are considered necessary for all FSNs. Others are optional.